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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classificati n ⁵: C09J 133/06, 7/02 D21C 5/02

A1

(11) International Publication Number:

WO 93/06184

(43) International Publication Date:

1 April 1993 (01.04.93)

(21) International Application Number:

PCT/US92/06731

(22) International Filing Date:

12 August 1992 (12.08.92)

(30) Priority data:

763,823

23 September 1991 (23.09.91) US

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(81) Designated States: AU, BR, CA, FI, JP, KR, European patent (AT, BE, CH, DE, DK, ES, FR, GB, GR, IE, IT, LU, MC, NL, SE).

Published

With international search report.

(54) Title: PRESSURE SENSITIVE ADHESIVE COMPOSITION WHICH IS REPULPABLE UNDER ACIDIC pH CONDITIONS

(57) Abstract

A pressure sensitive adhesive composition is described which is water-dispersible under acidic conditions. The composition comprises a terpolymer of a hydrophobic monomeric acrylic or methacrylic ester of a non-tertiary alcohol (monomer A); a polar monomer copolymerizable with said A monomer and having carboxylic, sulfonic acid or hydroxyl functionality or combinations thereof (monomer B); and a water soluble or water-dispersible macromolecular monomer of the formula X-Y-Z which is copolymerizable with A and B wherein X is a copolymerizable moiety containing a double bond, Y is a divalent linking group and Z is a water-dispersible polymeric moiety (monomer C). The combined acid groups employed in the B monomer are neutralized from 5 to 100 % based on the total amount of acid groups in B. The composition further comprises tackifier and/or plasticizers at a level such that the ratio of terpolymer to additives falls within a range of from 0.2:1 to 5:1. The pressure sensitive adhesive composition is dispersible under a variety ofpH conditions. Also described are tapes comprising the repulpable composition of the invention and their use in papermaking industry.

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PRESSURE SENSITIVE ADHESIVE COMPOSITION WHICH IS REPULPABLE UNDER ACIDIC pH CONDITIONS

5 Field of the invention

This invention relates to repulpable pressure sensitive adhesive compositions and tapes.

Background of the invention

Water-dispersible pressure sensitive adhesive compositions are used for many applications. For example they are used for fastening cloth on mammalian body coverings. Another field of use is papermaking and printing operations. There is required splicing of the end of one roll of paper to the beginning of another roll.

Examples of such water-dispersible pressure sensitive adhesives include U.S. Patent Nos. 3,865,770; 4,413,080 and 4,569,960 (Blake), U.S. Patent No. 20 3,441,430 (Peterson) and U.S. Patent No. 2,838,421 (Sohl). Blake discloses water-dispersible pressure sensitive adhesives for splicing carbonless paper which comprise a blend of acrylate: acrylic acid copolymers and systems which include tackifiers, plasticizers and 25 neutralizers. Peterson discloses aggressively tacky, water-soluble adhesives composed of a copolymer of a monocarboxylic acid and an alkoxy-alkyl ester, a plasticizer having at least one ether linkage, a cross linker and up to two parts of the hydroxide of a 30 monovalent metal ion. The hydroxide is used to neutralize sulfuric acid generated during the decomposition of a potassium persulfate catalyst used for copolymerization. Sohl discloses a mixture of water-soluble polyvinyl carboxylic acid and a 35 compatible hydroxy-polyalkylene permanent plasticizer which retains adequate tackiness and internal, cohesive strength even when exposed for a week or more at

humidities in the range of 0 to 90% relative humidity.

A variety of other pressure-sensitive adhesives are also said to be water-soluble or waterdispersible. For example U.S. Patent No. 4,413,082 (Gleichenhagen et al.) discloses a composition 5 comprising a copolymer of acrylic acid butyl ester and vinyl carboxylic acid and a plasticizer. The copolymer which contains free carboxyl groups is neutralized almost completely by the addition of potassium hydroxide preferably in excess. Gleichenhagen et al. 10 discloses that neutralization with potassium hydroxide normally brings about an excellent solubility of the pressure sensitive adhesive composition in water even at widely varying pH values. Solubility is measured in water without agitation. The only time reported to 15 achieve solubility lies between 5 and 12 minutes. adhesives disclosed by this reference are neutralized before being solubilized. However, in acidic media the salt resulting from neutralization will convert to the acid after a relatively brief time and the resulting 20 polymer will precipitate out of the solution. Materials being repulped often remain in the hydropulper for several hours. As a result, materials which will precipitate with time are not repulpable as described herein.

U.S. Patent No. 4,341,680 (Hauber et al.)
discloses a mixture of a copolymer of a monomeric ethyl
acrylate and a monomeric α,β- unsaturated aliphatic
monocarboxylic acid and a tertiary, ethoxylated n-alkyl
alkane diamine, where from 50% to 90% of the acid
groups of the monocarboxylic acid are neutralized.

European Patent Publication 0 352 442
discloses a pressure sensitive adhesive composition for
papermaking tapes. The composition is a water-soluble
contact adhesive based on a polyvinyl carbonic acid
with at least one softener containing a hydroxyl group
with a molecular weight below 1000 and curing agents
used to process paper.

U.S. Patent No. 3,152,940 (Abel et al.) discloses a pressure sensitive adhesive composition for photographic papermaking repulpable tapes. The preferred composition is an alkali-soluble copolymer of acrylic acid ester and acrylic acid and a water-soluble wax of hydroxy polyalkylene material which remains cohesive and tacky when exposed either to dry or humid atmospheric conditions.

U.S. Patent No. 4,338,432 (Eskay) discloses a pressure sensitive adhesive composition for making water-dispersible tapes for papermaking. The adhesive employs a copolymer of an acrylic acid compound, which is a homopolymer of acrylic acid or a copolymer of acrylic acid and esters of acrylic acid or methacrylic acid, a polyhydric alcohol plasticizer-tackifier, and a minor amount of monobasic caustic. Amounts of the base higher than 2.5 percent cause the adhesive to become elastic in nature rather than tacky.

Another water-soluble pressure sensitive

20 adhesive for papermaking tapes is disclosed in U.S.

Patent No. 3,096,202 (deGroot von Arx).

Polyvinylpyrrolidone, a polyfunctional monomeric crosslinking agent, a compatible plasticizer, and an activator catalyst to induce vinyl-type polymerization

25 are combined to produce a water-soluble, normally tacky pressure sensitive adhesive retaining its adhesiveness after subjection to elevated temperatures.

European Patent Publication 0 297 451
(Knutson et al.) discloses a hot melt adhesive
30 activated by moisture which is water-soluble and
alkali-dispersible to be recycled with paper products.
The hot melt adhesive comprises an N-acrylpolyakyleneimine and an acid functional compound.

Another water-soluble adhesive said to be
35 useful with paper products such as decals, labels, and
bumper stickers is disclosed in U.S. Patent No.
3,763,117 (McKenna et al.). The composition which has

aggressive tack and aqueous alkali removability comprises a hydroxy bearing monomer, a softening acrylate or methacrylate comonomer, and optionally, a hardening comonomer of either vinyl esters of alkanoic acids, ethyl or methyl acrylates, ethyl or methyl diesters of maleic or fumaric acids, acrylonitrile, methacrylonitrile, styrene, or vinyl chloride.

A transfer or splicing tape is disclosed in U.S. Patent No. 3,890,292 (Bohme et al.). The tape is formed with a water-dispersible adhesive composition of ionomeric copolymers of hydrophilic and hydrophobic monomers. The ionic monomer comprises a water-soluble alkali metal salt of an α,β - unsaturated monocarboxylic acid. The water-soluble non-ionic monomers are polyalkylene oxide condensation products. The composition maintains excellent tack characteristics at both high and low humidities and has improved moisture stability properties.

Yet another water-dispersible pressure 20 sensitive adhesive is disclosed in pending United States patent application 07/580,116 (Attorney docket number 45417USA8A) filed September 10, 1990. adhesive disclosed therein is formed from a blend of (a) 100 parts by weight of a copolymer of monomers of 25 (1) about 50-85 weight percent of at least one monomeric acrylic acid ester of nontertiary alkyl alcohol having 2-8 carbon atoms, and (2) about 50-15 weight percent of a vinyl carboxylic acid, (b) about 10 to about 250 parts by weight of a water-dispersible or 30 water-soluble plasticizer, (c) optionally from about 0 to about 100 parts by weight of an acrylate-compatible tackifier not irritating or toxic to mammalian skin; and (d) sufficient alkali metal hydroxide, if any, to neutralize from 0 to no more than 50 percent of acid 35 moieties in the copolymer, to cause the adhesive to retain pressure sensitive adhesive properties during

exposure to moisture and body fluids yet be dispersible in aqueous alkali solutions.

These pressure sensitive adhesives are soluble or dispersible under neutral or alkaline 5 conditions. However, they typically have poor watersolubility under acidic conditions. Moreover it is not believed that they are repulpable under acidic conditions as described herein. Accordingly a need exists for a pressure sensitive adhesive which is 10 repulpable (defined more fully hereinafter) under acidic conditions. For example, in papermaking and finishing operations, subsequent rolls are joined together with pressure sensitive adhesives (PSA) to form an endless web. The spliced areas are removed in 15 the mill or at the end user and recycled in the manufacturing process. The paper pulp obtained from these materials (scrap) must not be contaminated with insoluble sticky substances, originating from the PSA.

Such adhesives have to meet several

- 20 requirements in order to be useful:
 - In several processes the splice is subjected to elevated temperatures (200°C to 250°C) under high shear forces. The most severe conditions are typically observed in supercalender applications.
- 25 Any slippage, even fractions of millimeters could lead to blocking and subsequent web breakage.
 - A high cohesive strength is needed to withstand the high pressure/heat and resist sideways flowing (oozing) or penetration through the paper (bleeding).
 - Adequate repulpability under a variety of pH conditions including both alkaline and acidic conditions (but especially acidic conditions).
 - Good balanced adhesion properties
- 35 (tack/peel/shear)

30

- Useable with light weight coated (LWC) papers.

The present invention describes a pressure sensitive adhesive composition which possesses these properties. The adhesive composition is especially suitable for preparing pressure sensitive adhesive tapes which can be used in the paper industry for splicing papers. However, the adhesive can also be used for such other applications as labels, masking tapes, protective coverings, stamps and the like.

10 Summary of the invention

The present invention provides a pressure sensitive adhesive composition which has a pH-independent repulpability over a wide range of pH values. It has been discovered that a combination of (i) an acrylate copolymer grafted with an appropriate amount of water-dispersible segments, and (ii) selected additives provides an adhesive that is repulpable at a variety of pH values, particularly acidic pH values. The acrylate copolymer can be obtained using macromer technology. These segments help impart a pH-independent water-dispersibility to the formulation.

As used herein, the term "water-dispersible" means that the respective compositions are capable of being either dispersed in or dissolved in water alone at room temperature, such as 20°C, and atmospheric pressure.

The word "tackifier" is used herein to mean a compound which is added to the base adhesive polymer to impart or increase the ability of the adhesive to form a bond of measurable strength between two surfaces immediately upon contact of such surfaces with, at most, the application of low pressure at room temperature.

The word "plasticizer" is used herein to mean 35 a compound which reduces the Tg of the adhesive composition and results in a better flow and rapid wetting of the adherent.

The term "macromer" is used herein to mean a macromolecular monomer which is an oligomeric polymeric material with a number average molecular weight of preferably 300 to 30,000 and having a polymerizable 5 group.

The phrase "pressure sensitive adhesive" (PSA) is used herein to mean a composition which adheres two substrates together with, at most, the application of low pressure at room temperature.

The phrase "repulpable" is used herein to mean that the adhesive has an ERT rating (described hereinafter) of no more than 3.

The pressure sensitive adhesive composition of the present invention has several advantages. It is repulpable over a wide range a pH values as shown by reference to a European repulpability test described hereinafter. Additionally, the composition of the invention preferably shows balanced adhesive properties, like peel strength and heat resistance, as well as acceptable tack/adhesion to papers.

Tapes which utilize the pressure sensitive adhesive composition of the present invention may be provided as transfer tapes (e.g., no backing required), with a backing, etc. If a backing is employed, it may be covered on one or both sides with the pressure sensitive adhesive. The backing material may consist of a material which is repulpable under the same conditions as the pressure sensitive adhesive composition.

30

Detailed description

The pressure sensitive adhesive of the present invention comprises a composition which is water-dispersible segments and, therefore, repulpable over a pH range of from 2 to 11 when tested according to the European repulpability test described

hereinafter. Preferably the adhesive is repulpable in a pH range of from 2 to 7.

The composition utilizes, as one essential ingredient, a terpolymer formed by the copolymerization of A, B and C monomers. Additives, such as tackifiers and plasticizers are employed as a second essential ingredient. A variety of other materials may also be employed in the adhesive as will be more fully discussed hereinafter. However, these other materials are not essential to the adhesive.

The terpolymers useful in the invention preferably have an inherent viscosity in methyl ethyl ketone of from 0.4 to 3 g/dl. Most preferably the inherent viscosity is in the range of from 0.5 to 1.5 g/dl.

The A monomer useful in preparing the terpolymer is a hydrophobic monomeric acrylic or methacrylic ester of a non-tertiary alcohol, which alcohol contains from 2 to 14 carbon atoms and preferably from 4 to 12 carbon atoms. It is preferred that the non-tertiary alcohol can be an alkyl alcohol. The term "hydrophobic" is used herein to mean that the A monomer lacks substantial affinity for water, that is it neither substantially adsorbs nor absorbs water at room temperature.

Examples of monomers suitable for use as the A monomer include the esters of either acrylic acid or methacrylic acid with non-tertiary alcohols such as ethanol, 1-propanol, 2-propanol, 1-butanol, 2-butanol, 1-pentanol, 2-pentanol, 3-pentanol, 2-methyl-1-butanol, 1-hexanol, 2-hexanol, 2-methyl-1-pentanol, 3-methyl-1-pentanol, 2-ethyl-1-butanol, 3,5,5-trimethyl-1-hexanol, 3-heptanol, 1-octanol, 2-octanol, isooctylaclohol, 2-ethyl-1-hexanol, 1-decanol, 1-decanol, 1-dodecanol, 1-tridecanol, 1-tetradecanol and the like. The preferred A monomer is the ester of acrylic acid with butyl alcohol or isooctyl alcohol or a combination

thereof, although combinations of two or more different A monomers are suitable.

Monomer A is preferably present in an amount of 30 to 85% by weight based on the total monomer 5 content used to prepare the terpolymer. More preferably monomer A is present in an amount of 40 to 60% by weight. It has been found that the waterdispersibility of the terpolymer, and hence that of the adhesive, drops when isooctyl acrylate is employed at a 10 level of more than 50% by weight of the total monomers used to make the terpolymer.

The B monomer useful in preparing the terpolymer is polar and is copolymerizable with the A monomer. It has hydroxy, carboxy or sulfonic acid functionality or salts thereof. When a B monomer having a hydroxy functionality is employed, a second B monomer having an acidic group is also present.

Representative examples of monomers suitable for use as the B monomer are vinyl carboxylic acids

20 like acrylic acid, methacrylic acid, itaconic acid, maleic acid, fumaric acid, and B-carboxyethylacrylate.

Other monomers suitable as the B monomer include a vinyl sulfonic acid. Examples of such acids are 2-acrylamido-2-methylpropane sulfonic acid,

25 sulfoethyl methacrylate, sulfopropyl methacrylate, sodium styrene sulfonate, N-(3-sulfopropyl)-N-methacroyloxy-ethyl-N,N-dimethyl-ammonium betaine, 1-(3-sulfopropyl)-2-vinyl-pyridinium-betaine, bis-(3-sulfopropyl)-itaconic acid ester.

Yet other useful B monomers include 2hydroxyethylacrylate, 2-hydroxyethylmethacrylate,
hydroxypropylacrylate and hydroxypropylmethacrylate.
These each contain at least one hydroxy group and must
be combined with a B monomer having an acidic group.

The amount of monomer B should not exceed 50% by weight of total monomers. Monomer B is preferably present in an amount of from 2 to 50% by weight. More

preferably, monomer B is present in an amount of from 10 to 30% by weight based on total weight. Moreover, from 5 to 100% of the acidic groups of the B monomer are neutralized.

several neutralizing agents, such as amines and alkali metal hydroxides, can be used for neutralization of the acidic groups of the B monomer. The neutralization not only improves the repulpability/dispersibility of the adhesive, it also increases the cohesive strength of the adhesive composition. Preferably sodium and potassium hydroxide are used as neutralizing agents. Also preferred is the use of triethanolamine as co-neutralizing agent with sodium hydroxide or potassium hydroxide. The neutralizing agents can be regarded as ionic crosslinkers. The exact quantity of neutralizing agent employed is not critical.

The cohesive strength of the adhesive can also be adjusted with carefully selected and controlled 20 amounts of covalent crosslinkers to impart cohesiveness up to a point where the repulpability is not effected adversely. It may be desirable in some instances in the practice of the invention to increase internal strength or cohesiveness of the terpolymer by 25 crosslinking the terpolymer using conventional methods. Any of several well known chemical crosslinking agents may also be used. Crosslinkers as further additives to the pressure sensitive adhesive compositions can be bisamide, UCARTM 1100 (Union Carbide) and a polyamide epichlorhydrin compound. Typical crosslinking agents of the invention are those disclosed in US 4,418,120 (Kealy and Zink) and in US 4,569,960 (Blake).

The C monomer useful in preparing the terpolymer is a water-dispersible macromolecular monomer. Preferably the C monomer contains only one vinyl group copolymerizable with the A and B monomers.

5

Typically, the C monomer contains a plurality of hydrophilic sites which impart the required hydrophilicity of the monomer. Monomer C may be represented by the general formula I

 $X-Y-Z \tag{I}$

wherein X is a structural element comprising a moiety copolymerizable with the A and B monomers, Y is a divalent linking group, and Z is the hydrophilic site and comprises a water-dispersible polymeric moiety, i.e. one containing two or more monomer units which are essentially unreactive under free radical initiated copolymerizing conditions employed to form the pressure sensitive adhesive terpolymer.

A preferred X group of the C monomer is a 15 vinyl group of the general formula II

$$H_2C=CR^1-$$
 (II)

wherein R¹ is a hydrogen atom or a methyl group.

A preferred Y group of the C monomer is a divalent carbonyl group.

- A preferred Z moiety of the C monomer is an oligomeric/polymeric material with a molecular weight of 300 to 30,000. Preferably, the Z moiety is a polyethoxyloxazoline or a poly-n-vinylpyrrolidone or a polyalkyleneoxide structure. It is also possible to use a Z compound which is a copolymer of different compounds, e.g. a N-vinylpyrrolidone and acrylamide. If such a copolymer is prepared, one of the monomers, e.g. the arcylamide is present only in minor amounts, that is less than 10% by weight.
- C monomers of the type useful for the invention are not available commercially. One method of preparing C monomers uses commercially available preformed polymeric Z moieties and conventional chemical reactions for modifying them. For example, a mono-methoxypolyethlyeneglycolacrylate monomer can be prepared by esterification of a monohydroxyl-terminated poly(lower alkylene oxide) compound in an appropriate

solvent. To the solution is added a ∝,8-unsaturated carboxylic acid, preferably in presence of p-toluene sulfonic acid and refluxed for generally 16 to 18 hours. Excess acid is neutralized and the precipitate is filtered off.

Suitable monohydroxyl terminated poly(lower alkylene oxides) which may be used to prepare the C monomers using the above described procedure include Carbowax^M 750, Carbowax^M 550 and Carbowax^M 350.

addition reaction in which an isocyanate is reacted with a monohydroxyl or monoamine terminated poly(lower alkylene oxides) as well as other Z moieties as described above. For example a benzylisocyanate having a vinylic double bond in the side chain may be reacted with a monoamine terminated poly(lower alkylene oxide). The components are mixed and reacted for a few hours for quantitative conversion. A suitable monoamino terminated poly(lower alkylene oxide) is for example Jeffamine M 600 (Texaco).

The C monomer can be also prepared by first synthesizing a suitable Z moiety and then converting it

to a macromolecular monomer using known methods. For example, the Z moiety is may be formed by radical

25 polymerization of N-vinylpyrrolidone and acrylamide in the presence of a chain transfer agent for example amino-ethanethiol, using a radical chain initiator for example azobisisobutyronitrile in an appropriate organic solvent. Then, in a second step, the X and Y

moiety containing compound is reacted with the Z-moiety which bears primary amine groups as a result of the chain transfer reaction. A suitable reagent for introducing the X and Y groups is vinyl azlactone (available from SNPE, France). The vinyl-azlactone

35 ring can be opened yielding the C monomer.

Monomer C is preferably present in an amount of from 10 to 50% by weight, preferably in the amount of 20 to 50% by weight of the total monomers.

The terpolymer is combined with one or more

5 additive materials selected from the group consisting
of tackifiers and plasticizers to form a multicomponent
composition. Useful tackifiers have a glass transition
temperature higher than that of the base polymer
whereas useful plasticizers have a glass transition
10 temperature lower than that of the base polymer.

The tackifier is preferably selected from the group consisting of rosin or rosin derivatives, acidic rosin or rosin derivatives and/or acidic terpenephenolic resins. The plasticizer is preferably selected from the group consisting of polyoxyethylenealkyl-phenyletherphosphate, polyoxyethylenealkyletherphosphate, polyethylenealkyletherphosphate, polyethyleneglycolmonophenylether, octyl-phenoxypoly(ethyleneoxy)ethanol and/or nonylphenoxypoly(ethyleneoxy)ethanol.

The ratio of additives to terpolymer employed in the composition of the invention is in the range of 0.2:1 to 5:1. It is more preferred that the ratio be in the range of 0.5:1 to 1.5:1 and most preferred that the 25 ratio be 1:1.

A typical procedure for preparing the pressure sensitive adhesive composition of the present invention is as follows.

The monomers A, B and C are preferably

30 dissolved in an organic solvent, and copolymerized in
presence of a radical chain initiator, such as
azobisisobutyronitrile, and held at elevated
temperature, such as 45 to 80°C, for several hours.

The resulting polymer solution may then be 35 combined with the additives, e.g., the plasticizer and tackifier, and other adjuvants, e.g., the neutralizing agent and the crosslinker, if any. Alternatively, the

plasticizer may be added before the polymerization process. The blended pressure sensitive adhesive composition typically comprises from about 20% to 60% by weight in the solvent. The preferred organic solvent system comprises ethyl acetate, methanol, isopropanol and water.

Pressure sensitive adhesive tape of the invention may be made by applying the pressure sensitive adhesive composition to one or more surfaces of a substrate to form a tape. Preferably the substrate is a repulpable one such as a tissue paper. The adhesive tape may also be used as a transfer adhesive fortified with fibers for easy handling.

Release liners known to those skilled in the
15 art may cover and protect the exposed surfaces of the
pressure sensitive adhesive tape between manufacturing
and use. Commercially available release liners include
siliconized paper release liners.

20 <u>Examples</u>

The invention is further illustrated by the following nonlimiting examples.

The following test methods were used to obtain the data set forth in the examples.

25 A short description of certain test procedures will facilitate the evaluation of characteristics of the repulpable pressure sensitive adhesive composition or construction.

30 1. Adhesion To Paper.

This test measures 180° peel from paper after a dwell time of one minute. A 3.175 cm x 20 cm strip of 70 gram offset coated paper (BRO glossy, available from Scheufelen and Company, Lenningen, Germany) is laminated to a 5 cm x 30 cm stainless steel test panel using a double coated pressure sensitive adhesive tape. One end of a 2.54 cm x 30.0 cm strip of sample tape

having a 37.5 micron thick polyester backing is then laid over the offset paper. Uniform adhesive contact is assured by passing a 2 kg rubber roller over the sample at a speed of 30.5 cm/minute. The specimen is 5 tested within one minute. The tape is then doubled back on itself and peeled from the offset paper surface at 5.08 cm/minute using PSTC-3 (modified as described above) described in TEST METHODS for pressure sensitive tapes, Pressure Sensitive Tape Counsel, 9. edition, 10 Willmot Road, Suite 201, Deerfield, Illinois 60015-5165.

2. Repulpability

Two test methods are described. One is a qualitative method and one is a quantitative method. The quantitative test method described demonstrates a recently introduced procedure. In this test method, equipment and conditions are employed which try to duplicate hydropulper conditions, i.e. blending for approximately 10 minutes at 3,000 rpm. The adhesives are repulpable if they have a water-dispersibility rating of at most 3.

2(a). Water-dispersibility Test (D)
A layer of coated adhesive is subjected to
25 the following test. A few drops of water are sprinkled on the surface and the thus moistened surface rubbed with a finger. A water-dispersible material will become slippery and start to dissolve. This is a qualitative test which indicates the tendency of the 30 adhesive to be repulpable.

2(b). European Repulpability Test (ERT)

Both paper and adhesive tape are conditioned
at 23°C/50% relative humidity for 24 hours before

running the test. 0.3 g of double coated tape to be

35 tested is sandwiched between two strips of DIN A4

paper. Additional DIN A4 paper is added to provide a
total weight of 47.3 g in order to have a tape/paper

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ratio of 0.6%. Under certain circumstances, 4% tape versus paper can be employed to simulate extreme The samples are cut into 1 cm by 1 cm squares and introduced into a conventional blender with 5 2 l of demineralized water (23°C) resulting in a pulp content of 2.5% by weight. The pH is adjusted to the desired value with diluted sulfuric acid solution. disintegration process is run for 10 minutes as described in detail in a publication from the Verein 10 der Zellstoff und Papier-Chemiker und -Ingenieure (Society for Cellulose and Paper Chemists and Engineers) in their Merkblatt V/4/61 and V/6/61. stock is then diluted with tap water up to 10 1. sheets are produced from the pulp according to DIN 54 15 358 followed by a drying process of 10 minutes. produced sheets are visually inspected for fiber pulling upon removal of the resulting paper sheet from the substrate. Fiber pulling upon separation, and/or presence of speckles, is indicative of a residual 20 adhesive content Sheets are also inspected visually for adhesive nibs. After the drying operation, the top and bottom paper sheet are removed from the test sheet to observe any adhesive stringing between the test sheet and the top and/or bottom sheets. The top and/or 25 bottom sheets are then checked for damage (fibers torn The number, color and size of adhesive speckles per sheet is recorded.

Perfect repulpability implies no more than one small speckle (< 1 sq mm) on a sheet.

The results of the European repulpability test is rated according the following:

Acceptable

- 1 excellent No piece of undispersed adhesive.
- 2 very good One piece of undispersed adhesive less than 1 sq mm.
- 3 good Two pieces of undispersed adhesive less than 1 sq mm.

7

Not Acceptable

- 4 fair Three to five pieces of undispersed adhesive less than 1 sq mm.
 5 bad More than five pieces of undispersed
- adhesive less than 1 sq mm or at least one undispersed adhesive chip larger than 1 sq mm.
 - 6 very bad One or more original cuts of undispersed adhesive.
- 10 The resulting ERT rating reflects the worst of four measurements.

3. Heat-resistance

This test simulates the shear forces
15 encountered in a paper splice under elevated
temperature.

The end of a 3.17 cm strip of kraft paper or coated paper of approximately 20 cm length is covered with 2.54 cm of tape at right angles. The protective 20 liner is removed. A second strip of kraft paper is placed on top of the tape and directly in line with first strip of paper. The exposed tape is trimmed with a razor blade. The splice is rolled over once in each direction with 2 kg rubber covered roller. The test 25 specimen is then contacted to a curved heated surface maintained at 200°C. One end of the specimen is held in a clamp to prevent it from moving during the test. The other end of the specimen is threaded over a roll and allowed to hang free with a 1,250 gram weight 30 attached thereto so as to maintain the splice in contact with the heated surface. The time required for adhesive shear failure of the splice is then measured in seconds. The time is measured from the moment of first contact with the heated surface. This time is 35 recorded as heat-resistance of the splicing tape. the tape does not fail in 5 minutes, the amount of

slippage during the 5 minute period is recorded.

"C" monomer preparation

Polymeric vinyl terminated monomer identified in the claims and herein as "C" monomer is prepared as described below. The "C" monomers are identified in the forgoing descriptions and in the Tables as "C" monomers "C-1"-"C-4". Specific details of the X, Y and Z moieties are given in Table 1. In these preparations, the term "parts" means parts by weight unless otherwise noted.

10

Monomer "C-1"

A mono-methoxypolyethyleneglycolacrylate monomer with a molecular weight of 750 was prepared by introducing 100 parts of Carbowax™ 750 (Union Carbide) 15 and 100 parts of toluene to a flask which was equipped with a Dean-Stark condenser and a stirrer. solution was heated to reflux-temperature for two 11.3 parts of acrylic acid, 4.5 parts of ptoluene sulfonic acid, 3,000 ppm of Irganox™ PS 800 20 (Ciba-Geigy) and 500 ppm phenothiazine were added into this mixture. The solution was heated to refluxtemperature for another 16 hours. After cooling down to room temperature, excess acid was neutralized with 4.5 parts of calcium hydroxide. The formed precipitate 25 was filtered off. Subsequently the toluene was evaporated under reduced pressure to yield a solid 100% acrylate monomer.

Monomer "C-2"

To a vessel containing 25 parts by weight of m-isopropenyl-alpha, alpha-dimethyl benzylisocyanate (TMI -Cyanamid), 75 parts of Jeffamine M 600 (Texaco) were added and reacted for at least 2 hours to obtain a quantitative conversion.

Monomer "C-3"

97 parts of N-vinylpyrrolidone, 3 parts of acrylamide, 4 parts of aminoethanethiol, 2 parts of azobisisobutyronitrile, 80 parts of ethylacetate and 20 parts of methanol were added to a reaction vessel provided with a condenser, nitrogen inlet and stirrer. The solution was purged with nitrogen for 10 minutes and then was refluxed for 11 hours and cooled to room temperature. 10.7 parts of 2-methyl-4,410 dimethylvinylazlactone (SNPE, France) were then introduced into the solution and a quantitative ring opening reaction was established after 30 minutes.

Monomer "C-4"

dimethylformamide and 1.9 parts of methyltriflate were added to a polymerization bottle under nitrogen atmosphere. The polymerization reaction was carried out at 55°C for 75 hours at which point the reaction was terminated by the addition of 10 parts of N,N-dimethylaminoethylmethacrylate. After 30 minutes the reaction mixture was precipitated in 300 parts disopropylether. The ether layer was decanted and the residue dissolved in chloroform. This procedure was repeated twice. The presence of a poly-2-ethyloxazoline methacrylate monomer was identified by NMR-spectroscopy.

Monomer "C-5"

To a vessel containing 80 parts of Jeffamine™ M-600 (Texaco) was added 20 parts of 2-methyl-4,4-dimethyl- vinylazlactone. The combination was reacted at room temperature for one hour to obtain a quantative conversion.

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TABLE I "C"-Monomer

5	Desig- nation C-1	X-moiety H ₂ C=CH-	<u>Y-moiety</u> -C- 	<pre>Z-moiety -O-(polyethylene- oxide)</pre>
10	C-2	H ₂ C=C(CH ₃)-	O - N-C- H O	-N-(polypropylene- oxide) H
15	C-3	H ₂ C=CH-	-C-NC(CH ₃) ₂ -C- 	-N-(poly-N-vinyl- pyrrolidone) H
20	C-4	H ₂ C=C(CH ₃)-	-C-O(CH ₂) ₂ -N(CH ₃) ₂ -	-poly-2- ethyloxazoline
20	C-5	H ₂ C=CH-	-C-N-C(CH ₃) ₂ -C- 	-N (Polypropylene- oxide) H

Examples 1 - 12

The terpolymer was prepared by mixing the designated amounts of monomers A, B and C yielding 100 g of monomeric mixture. The monomers to be copolymerized were dissolved in ethylacetate in a one liter bottle. 0.3 weight percent of 2,2'-azobis-(2,4-dimethylvaleronitrile) as polymerization initiator and 10% by weight of isopropanol relative to the ethylacetate were added to the liter bottle. A nitrogen purge was introduced for two minutes at a flow rate of 1 liter/minute. The solution was held at 50°C for 24 hours, at which time the polymerization was 99% completed. To the solution was then added methanol and water to reduce viscosity and solids content.

To this solution of copolymer, the plasticizer,

40 tackifier and neutralizing agents were sequentially
blended with sufficient agitation to produce a clear
solution. After removal of the solvent the residue was
subjected to a repulpability test. The adhesive
compositions of the Examples 1 - 12 are given in Table

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II. The ERT rating, 180° peel and shear adhesion of the PSA of Table II are shown in Table IIa. Unless otherwise noted in Table II a, the ERT rating was measured at pH of 4.5.

of PSA
Composition
TABLE II

Neutr. Agent		NaOH/KOH/TEA 6 / 1 / 1	NaOH/KOH/TEA 6 / 1 / 1	NaOH	NaOH/KOH/TEA	6 / 1 / 1	NaOH	КОН	isooctylacrylate inherent viscosity of terpolymer arts terpolymer ts terpolymer						
Ves	Neutr.	20	20	09	09	45	45	45	09	45	75		100	100	acrylato viscos polymer lymer
Additives	₩ El	45 45	50 50	50 50	50 50	50 50	50 50	50 50	50 50	50 50	50 50		0 25	50 50	IOA = isooctylacrylate i.v. = inherent viscosi on 100 parts terpolymer 1100 parts terpolymer
	i.V.	0.7	2.24	ı	1.01	0.99	1.1	0.87	0.98	0.83	0.81		I	1.22	10A = i.v. = on 100 pc 100 pcr
	4 C-5		30												te i.v. = inherent visc in parts by weight based on 100 parts terpolym parts by weight based on 100 parts terpolymer ine
	C-3 C-4												20	- 10	y weigh weight
Terpolymer	C2										30		t	t	parts b rts by
Terpo	C-1	30	ı	25	30	20.	25	30	30	30	ı		ı	ı	מַסָּ בְּבַּ
ı	AA	20*	20	15	15	20	20	20	15	20	20		10	22	cry cry ac ize ize er
£	IOA	22.5	1	ı	1	ı	ı	1	55	20	ı		ı	ı	Notes to Table II BA = butyl acryla AA = acrylic acid P = plasticizer T = tackifier in TEA = triethanolam
•	x. BA	22.5	20	09	22	09	22	20	1	I,	10 50		11 70	12 68	Notes BA = AAA = PP = TT
	国	-	7	C	4	IJ	9	7	œ	Q	_			77	MHMHTT

Additionally 5 parts hydroxyethylacrylate as B monomer

The plasticizers (denoted by P in Table II) were:

Examples 1-2 Antarox™ CA 520 (GAF)
Examples 3-9 Gafac™ PE 510 (GAF)
Example 10 Antarox™ CA 520 (GAF)
5 Example 12 Gafac™ PE 510 (GAF)

The tackifier (denoted as T in Table II) was Foral™ AX in each Example where tackifier was present.

Table II a

10	Ex.	Repulp	ability	180° Pee	l <u>Heat</u>	Resistance
	_	ERT	<u>D</u>	(N/dm)	Shear(sec)	Slippage(mm)
	1	1	+	33.2	300+	< 0.25
	2	1*	•	25-27	300+	0.25
	3	1			300+	1.0
15	4	1	•	29.9	300+	0.5
	5	1		23.2	300+	0.5
	6	1		29.9	300+	< 0.5
	7	1		34.3	300+	0.25
	8	4		35.4	300+	< 0.5
2.0	9	. 1		33.2	300+	0.25
	10	1*	•	29.9	300+	< 0.25
	1 1		+	• .	300+	o
	12		+	٠.	300+	•
٠	*pH c	f 7			·	

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EXAMPLES 13-22

A terpolymer was prepared comprising BA/AA/C-1 (50/20/30 parts by weight) in each of the examples 13 to 21. The terpolymer of Example 22 had a composition of 55/15/30 parts by weight BA/AA/C-1. The terpolymers were prepared in the same manner as that of examples 1 to 12. The tackifier was ForalTM AX in an amount of 50 parts by weight. 50 parts by weight of the respective plasticizers was present. The neutralization was performed so that 75% of the acid functionality was neutralized with NaOH/KOH in a ratio of 3 to 1.

Table III illustrates the correlation between the molecular structure of the plasticizer and adhesion/heat-resistance properties. The evaluated plasticizers can be divided in two classes, i. e. ionic and nonionic. Each class can further be split into components with hydrophobic base aliphatic, aromaticaliphatic or aromatic groups. The names listed in the table are from top to bottom

GAFAC™ (GAF):

10 PE-510, RE-410, RM-410,
polyoxyethylenealkylphenylether-phosphate;
RB-400, RS-410, polyoxyethylenealkyletherphosphate
PYCAL 94 (ICI) polyethyleneglycol monophenylether
ANTAROXTM (GAF)

15 CA-520, CA-420, octylphenoxypoly(ethyleneoxy)ethanol

The GAF Antarox™ CA-520 plasticizer tends to generate the overall best properties. Also other types of low glass transition temperature water—soluble/dispersible materials can be employed as plasticizing agents, e.g. polyvinylmethylether (Lutonal™ - BASF), polyethyleneoxide, propyleneoxide block copolymers (Pluronic™ - BASF) and polyethylene and propylene glycols (Carbowax™ (Union Carbide)), Breox™ (BP Chemicals).

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TABLE III

Evaluation of Plasticizers

Transfer adhesives - 1 mil (0.0254 mm)

5	Ex.	<u>P</u>	<u>Heat r</u>	<u>esistance</u>	180°Peel <u>(N/dm)</u>
			<u>Shear</u>	Slippage	
		•	(sec)	(mm)	
-	13	PE-510	>300	<0.25	42.3
10	14	RE-410	>300	0.25	45.6
	15	RM-410	>300	<0.25	45.0
	16	RB-400	>300	0.25	15.7
	17	RS-410	>300	0.5	24.9
	18	PYCAL 94	>300	0.25	45.6
15	19	CA-520	>300	0.25	27.5
	20	CA-420		AF	
	21	CO-430	٠	AF	
	22	RM-410	>300	0.25	54.4
	_				•

P = Plasticizer

20 AF = adhesive failure

A rosin-based resin with an acid moiety, (Foral™ AX (Hercules)) was evaluated as a tackifier. Through neutralization of the acid function of the tackifier it 25 is possible to produce water-dispersible compounds. This class of tackifiers is suitable as acrylate-compatible resins.

On certain types of paper, insufficient initial adhesion was observed. This shortcoming was improved substantially by using alkanolamine such as triethanolamine as co-neutralizing agent. This is shown by Table IV.

EXAMPLES 23-27

A base polymer of IOA/BA/AA/C-1 (25/25/20/30 parts by weight) was prepared according to examples 1 to 12 using Foral[™] AX and Antarox[™] CA-520. 10 or 15 parts of acrylic acid were neutralized with sodium hydroxide,

potassium hydroxide and/or triethanolamine (TEA) in the ratios given in the parentheses of Table IV. The 180° peel adhesion (N/dm) were measured on double-coated tapes, each sample being coated with 31 grams per square meter on each side of the test specimen.

			Table IV	
	<u>Ex</u>	<pre>% Neutralization</pre>	Agent	180°Peel Adhesion (N/dm)
10			•	
	23	50	NaOH	12.2 - 15.5
	24	75	NaOH/KOH (3 / 1)	14.4 - 17.7
15	25	75	NaOH/TEA (3 / 1)	22.1 - 25.4
	26	75	NaOH/TEA (2 / 1)	33.2 - 36.5
	27	75	NaOH/KOH/TEA (6 / 1 / 1)	25.4 - 29.9

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Claims:

A pressure sensitive adhesive composition
which is repulpable at a pH of from 2 to 11 comprising
 (a) a terpolymer formed from A, B and C monomers
wherein,

A is a hydrophobic monomer selected from acrylic or a methacrylic ester of a non-tertiary alcohol, the alcohol having 2 to 14 carbon atoms, wherein A comprises from 30 to 85 % by total weight of the A plus B plus C;

B is a polar monomer copolymerizable with said A monomer and having hydroxyl, carboxyl, sulfonic acid functionality or combinations thereof, with the proviso that if B contains the hydroxyl functionality it is combined with a monomer having an acid group, wherein B comprises from 2 to 50 % by total weight of A plus B plus C, and wherein from 5 to 100% of any said acidic group has been neutralized; and

is a water-dispersible macromolecular monomer which is copolymerizable with A and B and has the formula X-Y-Z wherein X is a structural element comprising a moiety which is copolymerizable with A and B, Y is a divalent linking group joining X to Z, and Z is a water-dispersible moiety containing at least two units which are essentially unreactive under free radical initiated copolymerization conditions used to form the terpolymer, wherein C comprises from 10 to 50 % by total weight of A plus B plus C; and

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C

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(b) an additive selected from the group consisting of tackifiers, plasticizers, and combinations thereof wherein the ratio of said additive to said terpolymer is in the range of from 0.2:1 to 5:1.

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2. The adhesive according to claim 1 where the A monomer is selected from said acrylic or methacrylic ester of said non-tertiary alcohol.

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- 3. The adhesive according to claim 1 wherein the non-tertiary alcohol is selected from butyl and isooctyl alcohol.
- 4. The adhesive according to claim 1 wherein the B monomer is a vinyl carboxylic acid selected from acrylic acid, methacrylic acid, itaconic acid, maleic acid, fumaric acid, β-carboxyethylacrylate, salts of these monomeric acids and combinations thereof.

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5. The adhesive according to claim 1 where the polar B monomer contains at least one vinyl sulfonic acid functionality selected from 2-acrylamido-2-methylpropane sulfonic acid, sulfoethyl methacrylate, sulfopropyl methacrylate, sodium styrene sulfonate, N-(3-sulfopropyl)-N-methacroyloxyethyl-N,N-dimethyl-ammonium betaine, 1-(3-sulfopropyl)-2-vinylpyridinium-betaine, bis-(3-sulfo-propyl)-itaconic acid ester, salts of these monomers, and combinations thereof.

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6. The adhesive according to claim 1 wherein the B monomer is a combination of (i) one or more hydroxy functional materials selected from hydroxyethylacrylate, hydroxypropyl-acrylate, 135 hydroxyethylmethacrylate and hydroxypropyl-methacrylate and (ii) and acidic group containing material selected

from acrylic acid, methacrylic acid and an acid with vinyl sulfonic functionality.

- 7. The adhesive according to claim 1 where the C 5 monomer has a molecular weight of 300 to 30,000.
- 8. The adhesive according to claim 1 wherein the Z portion of the C monomer is selected from polyethyloxazoline, N-vinylpyrrolidone-acrylamide 10 copolymer, and a polyalkyleneoxide copolymer, and combinations thereof.
- 9. The adhesive of claim 1 wherein said tackifier has a glass transition temperature higher than said terpolymer and said plasticizer has a glass transition temperature lower than said terpolymer.
- 10. The adhesive of claim 9 where the tackifier is selected from the group of rosin or rosin 20 derivatives, acidic rosin or rosin derivatives, acidic terpenephenolic resins, and combinations thereof.
- 11. The adhesive of claim 9 where the plasticizer is selected from the group of polyoxyethylenealkyl-phenyletherphosphate, polyoxyethylenealkyletherphosphate, polyethyleneglycolmonophenylether, octyl-phenoxypoly(ethyleneoxy)ethanol, nonylphenoxypoly(ethyleneoxy)ethanol, and combinations thereof.
- 30 12. A crosslinked adhesive according to claim 1.
- 13. The adhesive of claim 1 further comprising a neutralizing agent selected from alkanol amines, water-soluble/dispersible alkali metal hydroxides and 35 combinations thereof.

- 14. The adhesive of claim 1 wherein said terpolymer has an inherent viscosity of from 0.4 to 3.0.
- 15. Process for the preparation of a pressure sensitive adhesive of claim 1, comprising the steps of:

copolymerizing the monomers A, B and C in presence of a radical chain initiator,

adding to the formed copolymer, a plasticizer, tackifier, a neutralizing agent and optionally, a cross-linking agent, in an organic solvent system comprising ethyl acetate, methanol, isopropanol and/or water.

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- 16. Pressure sensitive adhesive tape comprising the composition of claim 1.
- 17. Pressure sensitive adhesive tape of claim 16
 20 having at least one pressure sensitive adhesive
 composition of claim 1 on a backing layer.
 - 18. Pressure sensitive adhesive tape of claim 16 being a transfer or splicing tape for paper industries.

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- 19. A substrate bearing a layer of a pressure sensitive adhesive tape according to claim 1.
- 20. A repulpable splice comprising at least two sections of paper joined together by a pressure sensitive adhesive according to claim 1.

INTERNATIONAL SEARCH REPORT

International Application No

PCT/US 92/06731

I. CLASSIFICATION OF SUBJ	ECT MATTER (If several classificati	on symbols apply, indicate all) ⁶	
According to International Paten Int.Cl. 5 C09J133/	t Classification (IPC) or to both Nation 06; C09J7/02;	al Classification and IPC D21C5/02	
II. FIELDS SEARCHED	·		
•	Minimum Do	cumentation Searched?	
Classification System	I	Classification Symbols	
Int.C1. 5	CO9J ; CO8F		
	Documentation Searched of to the Extent that such Documentation	other than Minimum Documentation ents are Included in the Fields Searched [®]	
III. DOCUMENTS CONSIDER			T To Lawrence Clarine No.13
Category ° Citation of D	ocument, 11 with indication, where app	ropriate, of the relevant passages 12	Relevant to Claim No.13
MANUFAC 1 June see pag	e 4, line 16 - page 6 examples 102-112; ta	6, line 27;	1-8,10, 12,14-19
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considered to be of parti "E" earlier document but put filling date "L" document which may thr which is cited to establis citation or other special "O" document referring to an other means "P" document published prio- later than the priority da IV. CERTIFICATION Date of the Actual Completion of	eneral state of the art which is not cular relevance elished on or after the international ow doubts on priority claim(s) or the publication date of another reason (as specified) a oral disclosure, use, exhibition or to the international filing date but the claimed	"I" later document published after the inters or priority date and not in conflict with cited to understand the principle or theo invention "X" document of particular relevance; the clicanot be considered novel or cannot be involve an inventive step "Y" document of particular relevance; the clicanot be considered to involve an inventive document is combined with one or more ments, such combination being obvious in the art. "A" document member of the same patent factors of Mailing of this International Sec. 2.5. 11. 92	almed invention considered to timed invention considered to timed invention tive step when the other such docuton a person skilled mily
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ANNEX TO THE INTERNATIONAL SEARCH REPORT ON INTERNATIONAL PATENT APPLICATION NO. US 9206731 63494

This annex lists the patent family members relating to the patent documents cited in the above-mentioned international search report. The members are as contained in the European Patent Office EDP file on The European Patent Office is in no way liable for these particulars which are merely given for the purpose of information. 17/11/92

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